

wherein said Type I IFN has a sequence consisting essentially of the sequence of

- a) a native Type I IFN;
- b) a fragment of a) which has Type I IFN receptor agonist or antagonist activity;
- c) a variant of a) or b) which has at least 70% sequence identity with a) or b) and which has Type I IFN receptor agonist or antagonist activity; or
- d) a variant of a) or b) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding a) or b) under moderately stringent conditions and which has Type I IFN receptor agonist or antagonist activity;

C1  
Cont or a salt or functional derivative of a), b), c), or d) which has Type I IFN receptor agonist or antagonist activity; and

wherein said IFNAR has a sequence consisting essentially of the sequence of

- e) a native human IFNAR polypeptide chain;
- f) a fragment of e) which has IFNAR receptor agonist or antagonist activity;
- g) a variant of e) or f) which has at least 70% sequence identity with e) or f) and which has IFNAR receptor agonist or antagonist activity;

h) a variant of e) or f) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding e) or f) under moderately stringent conditions and which has IFNAR biological activity;

C<sup>1</sup>  
cont or a salt or functional derivative of e), f), g), or h) which has IFNAR biological activity,

with the proviso that when said Type I IFN and said IFNAR are administered separately and said complex is formed *in vivo*, the amount of IFNAR administered is an amount effective to prolong the *in vivo* effect of the Type I IFN.

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13 10 (Twice-amended). An isolated molecule comprising a complex of a Type I interferon (IFN) and a subunit of the human interferon  $\alpha/\beta$  receptor (IFNAR) which is capable of binding to the Type I IFN of the complex, in which said Type I IFN is bound to said IFNAR by a covalent bond or a peptide bond,

C<sup>2</sup> wherein said Type I IFN has a sequence consisting essentially of the sequence of

- a) a native Type I IFN;
- b) a fragment of a) which has Type I IFN receptor agonist or antagonist activity;

c) a variant of a) or b) which has at least 70% sequence identity with a) or b) and which has Type I IFN receptor agonist or antagonist activity; or  
d) a variant of a) or b) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding a) or b) under moderately stringent conditions and which has Type I IFN receptor agonist or antagonist activity;  
or a functional derivative of a), b), c), or d) which has Type I IFN receptor agonist or antagonist activity; and  
wherein said IFNAR has a sequence consisting essentially of the sequence of

- C<sup>2</sup>  
cont
- e) a native human IFNAR polypeptide chain;
  - f) a fragment of e) which has IFNAR biological activity;
  - g) a variant of e) or f) which has at least 70% sequence identity with e) or f) and which has IFNAR biological activity; or
  - h) a variant of e) or f) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding e) or f) under moderately stringent conditions and which has IFNAR biological activity;

C2  
cont or a salt or functional derivative of e), f), g), or h) which has IFNAR biological activity.

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27 22 (Amended). A pharmaceutical composition consisting essentially of a pharmaceutically acceptable carrier and a complex of a Type I interferon (IFN) and a subunit of the human interferon  $\alpha/\beta$  receptor (IFNAR) which is capable of binding to the type I IFN of the complex,

wherein said Type I IFN has a sequence consisting essentially of the sequence of

- C3
- a) a native Type I IFN;
  - b) a fragment of a) which has Type I IFN receptor agonist or antagonist activity;
  - c) a variant of a) or b) which has at least 70% sequence identity with a) or b) and which has Type I IFN receptor agonist or antagonist activity; or
  - d) a variant of a) or b) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding a) or b) under moderately stringent conditions and which has Type I IFN receptor agonist or antagonist activity;
- or a salt or functional derivative of a), b), c), or d) which has Type I IFN receptor agonist or antagonist activity; and
- wherein said IFNAR has a sequence consisting essentially of the sequence of
- e) a native human IFNAR polypeptide chain;

f) a fragment of e) which has IFNAR receptor agonist or antagonist activity;

g) a variant of e) or f) which has at least 70% sequence identity with e) or f) and which has IFNAR receptor agonist or antagonist activity; or

h) a variant of e) or f) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding e) or f) under moderately stringent conditions and which has IFNAR receptor agonist or antagonist activity;

or a salt or functional derivative of e), f), g), or h) which has IFNAR biological activity.

*C<sup>3</sup> Cont.*

12 ~~23~~ (Amended). A method for potentiating the biological effects of Type I interferon (IFN), comprising:

administering to a patient in need of Type I IFN therapy a subunit of the human interferon  $\alpha/\beta$  receptor (IFNAR) which is capable of binding to the Type I IFN to be potentiated, in an amount effective to provide such IFN therapy,

wherein said IFNAR has a sequence consisting essentially of the sequence of

- a) a native human IFNAR polypeptide chain;
- b) a fragment of a) which has IFNAR receptor agonist or antagonist activity;

- c) a variant of a) or b) which has at least 70% sequence identity with a) or b) and which has IFNAR receptor agonist or antagonist activity; or
- d) a variant of a) or b) which is encoded by a DNA sequence which hybridizes to the complement of the native DNA sequence encoding a) or b) under moderately stringent conditions and which has Type I IFN receptor agonist or antagonist activity;

or a salt or functional derivative of a), b), c), or d) which has IFNAR receptor agonist or antagonist activity.

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<sup>C3</sup>  
cont  
10 ~~25~~ (Amended). A method in accordance with claim 1, wherein said native human IFNAR polypeptide chain of e) is the extracellular domain of a native human IFNAR polypeptide chain.

<sup>C4</sup>  
13 ~~21~~ ~~26~~ (Amended). A molecule in accordance with claim ~~10~~, wherein said native human IFNAR polypeptide chain of e) is the extracellular domain of a native human IFNAR polypeptide chain.

~~28~~ ~~27~~ (Amended). A pharmaceutical composition in accordance with claim <sup>27</sup>~~22~~, wherein said native human IFNAR polypeptide chain of e) is the extracellular domain of a native human IFNAR polypeptide chain.

11 ~~28~~ (Amended). A method in accordance with claim 3, wherein said native human IFNAR polypeptide chain of e) is the

C4  
cont extracellular domain of a native human IFNAR polypeptide  
chain.

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